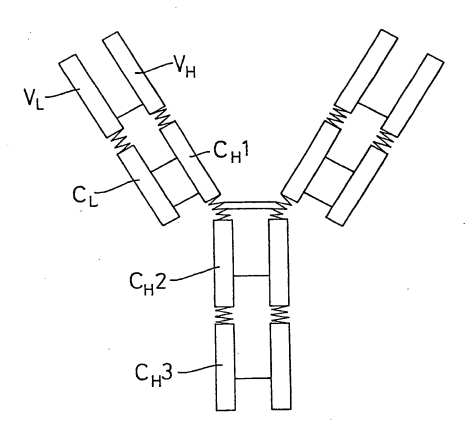
Fig. 1

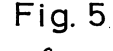


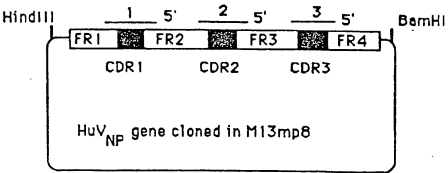
domains inter-domain sections Mdisulphide bonds ٧ variable constant C light chain heavy chain

Н

31 NOVYT SYUNH	SO CDR2 65 YUFYHGTSDOTTPLRS RIDPNSGGTKYNEKFKS	24 95 CDR3 102 R NLIRGCIDU R YDYYGSSYFDY	Fig. 2
YVOLQESGPGLURPSQTLSLTCTUSGSTFS QUQLQQPGRELUKPGRSUKLSCKRSGYTFT	36 HUROPPGRGLEHIG HUKORPGRGLEHIG	66 RUTHLUDISKHQFSLRLSSUTARDTAUYYCAR KATLTUDKPSSTAYHQLSSLTSEDSRUYYCAR	103 HGQGSLUTUSS 113 HGQGTTLTUSS
EUM 31-8	NELM 81-8	NEUM B1-8	NELT B1-8

HindIII -48	-23	-7
· .*ATGCARATCCTCTGAATC	TACATGGTAAATATAC	GTTTGTCTATAC
RNA starts RN CACAAACAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	IA starts TCTACAGTTACTGAGG	CACACAGGACCTC
NP leader		Splice
M G M S C I I L F L ACCATGGGATGGAGCTGTATCATCCTCTTCTT		
ACAGTAGCAGGCTTGAGGTCTGGACATATATA Splice, 1		10
	LQLESG	PGLUR
TCTCTCCACAGGTGTCCACTCCCAGGTCCAAC	TGCA GGAGAGCGGTO	CRGGTCTTGTGRG
	5.1	
15 20 P S Q T L S L T C T	25 U S G S T	30 CDR1
#CCT##CC##############################	הדהדרדההרפהרפררי	TEREFORETRE
4 1 6	5	8-7-
	45	50 CDR2 52R
M H H U R Q P P G R	GLEHI	GRIDP
GRTGCACTGGGTGAGACAGCCACCTGGACGA		
7	- 10b	12/14
55 CDR2 60		70
N S G G T K Y N E K TRATAGTGGTGGTACTARGTACARTGAGRAG		
11-12/14-13a-13a-12/14	13b	J 15
75 80 8		
TSKNQFSLRL	SSUTA	A D T. A V
CACCAGCAAGAACCAGTTCAGCCTGAGACTC	RGCRGCGTGRCRGCCC 7	CCGRCACCGCGGT
15		20
90 95 CDR3		105
Y Y C A R Y D Y Y G CTATTATTGTGCRAGATACGATTACTACGGT		
1921		.23
		26a
SPICE SLUTUSS CRECTCETCACAGT	BamH	I
	· 193bp · · · · 3'	
	-	ia. 3





D1.3 CDR1 oligonucleotide 5' CTG,TCT,CAC,CCA,GTT,TAC,ACC,ATA,GCC,GCT,GAA,GGT,GCT

FR2

D1:3 CDR1

FR1

D1.3 CDR2 oligonucleotide
5' CAT,TGT,CAC,TCT,GGA,TTT,GAG,AGC,TGA,ATT,ATA,GTC,TGT,

FR3

D1.3 CDR2

GTT,TCC,ATC,ACC,CCA,AAT,CAT,TCC,AAT,CCA,CTC '

D1.3 CDR2

FR2

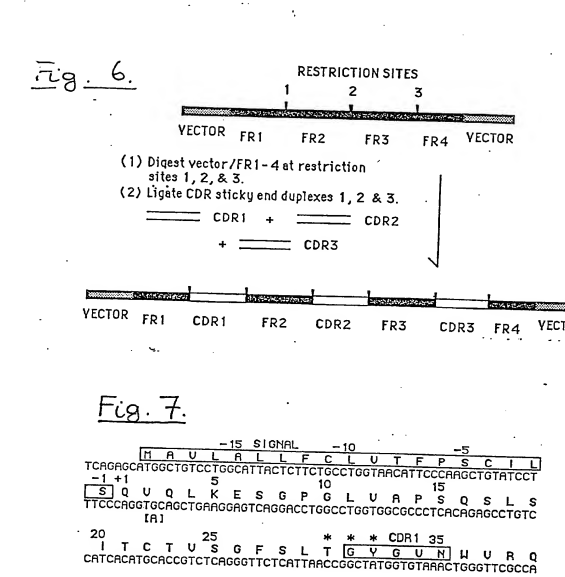
D1.3 CDR3 oligonucleotide
5' GCC,TTG,ACC,CCA,GTA,GTC,AAG,CCT,ATA,ATC,TCT,CTC,TCT,

FR4

D1.3 CDR3

TGC,ACA,ATA

FR3



40 45 50 * * * 55 CDR2
PPGKGLEWLGMINGGDGNTGGAACACAGACTA

TAATTCAGCTCTCAAATCCAGACTGAGCATCAGCAAGGACAACTCCAAGAGCCAAGTTTT

105 DVRLDYWGQGTTLTVSS AGATTATAGGCTTGACTACTGGGGCCAAGGCACCACTCTCACAGTCTCCTCA

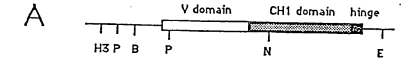
CDR3

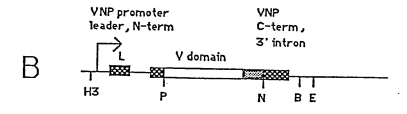
LKMNSLHTDDTARYYCARER сттянанатонаснототосносностоятся спостоя спосто

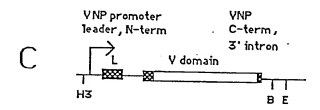
65 70 75 KSRLSISKDNSKSQUF

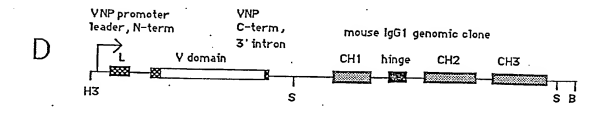
Fig. 8

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H3 = HindIII, P = Pstl, B = BamH1, N = Ncol, E = EcoR1, H2 = HindII

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DNA techniques.

However, colleagues of the present Inventor have devised a process whereby chimeric antibodies in which both parts of the protein are functional can be secreted. The process, which is disclosed in International Patent Application No. PCT/GB85/00392 (Neuberger et al. and Celltech Limited), comprises:

a) preparing a replicable expression vector including a suitable promoter operably linked to a DNA sequence comprising a first part which encodes at least the variable domain of the heavy or light chain of an Ig molecule and a second part which encodes at least part of a second protein;

 b) if necessary, preparing a replicable expression vector including a suitable promoter operably linked to a DNA sequence which encodes at least the variable domain of a complementary light or heavy chain re spectively of an Ig molecule;

c) transforming an immortalised mammalian cell line with the or both prepared vectors; and

d) culturing said transformed cell line to produce a chimeric antibody.

The second part of the DNA sequence may encode:

i) at least part, for instance the constant domain of a heavy chain, of an lg molecule of different species, 15 class or subclass;

ii) at least the active portion or all of an enzyme;

iii) a protein having a known binding specificity;

iv) a protein expressed by a known gene but whose sequence, function or antigenicity is not known; or v) a protein toxin, such as ricin.

20 The above Neuberger application only shows the production of a chimeric antibodies in which complete variable domains are coded for by the first part of the DNA sequence. It does not shown any chimeric antibodies in which the sequence of the variable domain has been altered.

The present invention, in a first aspect, provides an altered antibody in which at least parts of the CDRs in the light or heavy chain variable domains have been replaced by analogous parts of CDRs from an antibody of different specificity.

The determination as to what constitutes a CDR and what constitutes a framework region was made on the basis of the amino-acid sequences of a number of Igs. However, from the three dimensional structure of a number of Igs it is apparent that the antigen binding site of an Ig variable domain comprises three looped regions supported on sheet-like structures. The loop regions do not correspond exactly to the CDRs, 30 although in general there is considerable overlap.

Moreover, not all of the amino-acid residues in the loop regions are solvent accessible and in one case, amino-acid residues in the framework regions are involved in antigen binding. (Amit, A.G., Mariuzza, R.A., Phillips, S.E.V. and Poljak, R.J., Science, 233, 747-753, 1986).

It is also known that the variable regions of the two parts of an antigen binding site are held in the correct 35 orientation by inter-chain non-covalent interactions. These may involve amino-acid residues within the CDRs.

Thus, in order to transfer the antigen binding capacity of one variable domain to another, it may not be necessary to replace all of the CDRs with the complete CDRs from the donor variable region. It may be necessary only to transfer those residues which are accessible from the antigen binding site, and this may 40 involve transferring framework region residues as well as CDR residues.

It may also be necessary to ensure that residues essential for inter-chain interactions are preserved in the acceptor variable domain.

Within a domain, the packing together and orientation of the two disulphide bonded β-sheets (and therefore the ends of the CDR loops) are relatively conserved. However, small shifts in packing and orientation of these β-sheet do occur (Lesk, A.M. and Chothia, C., J. Mol. Biol., 160, 325-342, 1982). However, the packing together and orientation of heavy and light chain variable domains is relatively conserved (Chothio, C., Novotny, J., Bruccoleri, R. and Karplus, M., J. Mol. Biol., 186, 651-653, 1985). These points will need to be borne in mind when constructing a new antigen biding site so as to ensure that packing and orientation are not altered to the deteriment of antigen binding capacity.

50 It is thus clear that merely by replacing one or more CDRs with complementary CDRs may not always result in a functional altered antibody. However, given the explanations set out above, it will be well within the competence of the main skilled in the art, either by carrying out routine experimentation or by trial and error testing to obtain a functional altered antibody.

Preferably, the variable domains in both the heavy and light chains have been altered by at least partial 55 CDR replacement and, if necessary, by partial framework region replacement and sequence changing. Although the CDRs may be derived from an antibody of the same class or even subclass as the antibody from which the framework regions are derived, it is envisaged that the CDRs will be derived from an antibody of different class and preferably from an antibody from a different species.

Thus, it is envisaged, for instance, that the CDRs from a mouse antibody could be grafted onto the frame-60 work regions of a human antibody. This arrangement will be of particular use in the therapeutic use of monoclonal antibodies.

At present, when a mouse monoclonal antibody or even a chimeric antibody comprising a complete mouse variable domain is injected into a human, the human body's immune system recognises the mouse variable domain as foreign and produces an immune response thereto. Thus, on subsequent injections of the mouse antibody or chimeric antibody into the human, its effectiveness is considerably reduced by the action

of the body's immune system against the foreign antibody. In the altered antibody of the present invention, only the CDRs of the antibody will be foreign to the body, and this should minimise side effects if used for human therapy. Although, for example, human and mouse framework regions have characteristic features which distinguish human from mouse CDRs. Thus, an antibody comprised of mouse CDRs in a human frame-5 work may well be no more foreign to the body than a genuine human antibody. 5 Even with the altered antibodies of the present invention, there is likely to be an anti-idiotypic response by the recipient of the altered antibody. This response is directed to the anitbody binding region of the altered antibody, It is believed that at least some anti-idiotype antibodies are directed at sites bridging the CDRs and the framework regions. It would therefore be possible to provide a panel of antibodies having the same 10 partial or complete CDR replacements but on a series of different framework regions. Thus, once a first 10 altered antibody became therapeutically ineffective, due to an anti-idiotype response, a second altered antibody from the series could be used, and so on, to overcome the effect of the anti-idiotype response. Thus, the useful life of the antigen-binding capacity of the altered antibodies could be extended. Preferably, the altered antibody has the structure of a natural antibody or a fragment thereof. Thus, the 15 altered antibody may comprise a complete antibody, an (Fab')₂ fragment, an Fab fragment, a light chain 15 dimer or a heavy chain dimer. Alternatively, the altered antibody may be a chimeric antibody of the type described in the Neuberger application referred to above. The production of such an altered chimeric antibody can be carried out using the methods described below used in conjunction with the methods described in the Neuberger application. The present invention, in a second aspect, comprises a method for producing such an altered antibody 20 comprising; a) preparing a first replicable expression vector including a suitable promoter operably linked to a DNA sequence which encodes at least a variable domain of an Ig heavy or light chain, the variable domain comprising framework regions from a first antibody and CDRs comprising at least parts of the CDRs from a 25 second antibody of different specificity; 25 b) if necessary, preparing a second replicable expression vector including a suitable promoter operably linked to a DNA sequence which encodes at least the variable domain of a complementary Ig light or heavy chain respectively; c) transforming a cell line with the first or both prepared vectors; and d) culturing said transformed cell line to produce said altered antibody. 30 The present invention also includes vectors used to transform the cell line, vectors used in producing the transforming vectors, cell lines transformed with the transforming vectors, cell lines tranformed with preparative vectors, and methods for their production. Preferably, the cell line which is transformed to produce the altered antibody is an immortalised mam-35 malian cell line, which is advantageously of lymphoid origin, such as a myeloma, hybridoma, trioma or 35 quadroma cell line. The cell line may also comprise a normal lymphoid cell, such as a B-cell, which has been immortalised by transformation with a virus, such as the Epstein-Barr virus. Most preferably, the immortalised cell line is a myeloma cell line or a derivative thereof. Although the cell line used to produce the altered antibody is preferably a mammalian cell line, any other 40 suitable cell line, such as a bacterial cell line or a yeast cell line, may alternatively be used. In particular, it is 40 envisaged that E. Coli derived bacterial strains could be used. It is known that some immortalised lymphoid cell lines, such as myeloma cell lines, in their normal state secrete isolated Ig light or heavy chains. If such a cell line is transformed with the vector prepared in step a) of the process of the invention, it will not be necessary to carry out step b) of the process, provided that the 45 45 normally secreted chain is complementary to the variable domain of the lg chain encoded by the vector However, where the immortalised cell line does not secrete or does not secrete a complementary chain, it will be necessary to carry out step b). This step may be carried out by further manipulating the vector produced in step a) so that this vector encodes not only the variable domain of an altered antibody light or heavy 50 50 chain, but also the complementary variable domain. Alternatively, step b) is carried out by preparing a second vector which is used to transform the immortalised cell line. This alternative leads to easier construct preparation, but may be less preferred than the first alternative in that it may not lead to as efficient production of antibody. The techniques by which such vectors can be produced and used to transform the immortalised cell lines 55 are well known in the art, and do not form any part of the invention. 55 In the case where the immortalised cell line secretes a complementary light or heavy chain, the transformed cell line may be produced for example by transforming a suitabe bacterial cell with the vector and then fusing the bacterial cell with the immortalised cell line by spheroplast fusion. Alternatively, the DNA may be directly introduced into the immortalised cell line by electroporation. The DNA sequence encoding 60 60 the altered variable domain may be prepared by oligonucdeotide synthesis. This requires that at least the framework region sequence of the acceptor antibody and at least the CDRs sequences of the donor antibody are known or can be readily determined. Although determining these sequences, the synthesis of the DNA from oligonucleotides and the preparation of suitable vectors is to some extent laborious, it involves the use of known techniques which can readily be carried out by a person skilled in the art in light of the teaching 65 65 given here.

If it was desired to repeat this strategy to insert a different antigen binding site, it would only require the synthesis of oligonucleotides encoding the CDRs, as the framework oligonucleotides can be re-used. A convenient variant of this technique would involve making a symthetic gene lacking the CDRs in which the four framework regions are fused together with suitable restriction sites at the junctions. Double stranded 5 synthetic CDR cassettes with sticky ends could then be ligated at the junctions of the framework regions. A 5 protocol for achieving this variant is shown diagrammatically in Figure 6 of the accompanying drawings. Alternatively, the DNA sequence encoding the altered variable domain may be prepared by primer directed oligonucleotide site-directed mutagenesis. This technique in essence involves hybridising an oligonucleotide coding for a desired mutation with a single strand of DNA containing the region to be mutated 10 and using the single strand as a template for extension of the oligonulcleotide to produce a strand containing 10 the mutation. This technique, in various forms, is described by : Zoller, M.J. and Smith, M., Nuc. Acids Res., 10, 6487-6500, 1982; Norris, K., Norris F., Christiansen, L. and Fiil, N., Nuc. Acids Res., 11, 5103-5112, 1983; Zoller, M.J. and Smith, M., DNA, 3, 479-488 (1984); Kramer, W., Schughart, K. and Fritz, W.-J., Nuc. Acids Res., 10, 6475-6485, 1982, For various reasons, this technique in its simplest form does not always produce a high frequency of 15 mutation. An improved technique for introducing both single and multiple mutations in an M13 based vector, has been described by Carter et al. (Carter, P., Bedouelle H. and Winter, G., Nuc, Acids Res., 13, 4431-4443, Using a oligonucleotide, it has proved possible to introduce many changes simultaneously (as in Carter et 20 al., loc. cit.) and thus single oligonucleotides, each encoding a CDR, can be used to introduce the three CDRs 20 from a second antibody into the framework regions of a first antibody. Not only is this technique less laborious than total gene synthesis, but it represents a particularly convenient way of expressing a variable domain of required specificity, as it can be simpler than talloring an entire V_H domain for insertion into an expression plasmid. The oligonucleotides used for site-directed mutagenesis may be prepared by oligonucleotide synthesis or 25 may be isolated from DNA coding for the variable domain of the second antibody by use of suitable restriction enzymes. Such long oligonucleotides will generally be at least 30 bases long and may be up to over 80 bases in length. The techniques set out above may also be used, where necessary, to produce the vector of part (b) of the 30 30 process. The method of the present invention is envisaged as being of particular use in "humanising" non-human monoclonal antibodies. Thus, for instance, a mouse monoclonal antibody against a particular human cancer cell may be produced by techniques well known in the art. The CDRs from the mouse monoclonal antibody may then be partially or totally grafted into the framework regions of a human monoclonal antibody, which is 35 then produced in quantity by a suitable cell line. The product is thus a specifically targetted, essentially 35 human antibody which will recognise the cancer cells, but will not itself be recognised to any significant degree, by a human's immune system, until the anti-idiotype response eventually becomes apparent. Thus, the method and product of the present invention will be of particular use in the clinical environment. The present invention is now described, by way of example only, with reference to the accompanying 40 drawings, in which: 40 Figure 1 is a schematic diagram showing the structure of an IgG molecule; Figure 2 shows the amino acid sequence of the V_H domain of NEWM in comparison with the V_H domain of Figure 3 shows the amino acid and nucleotide sequence of the HuV_{NP} gene; 45 Figure 4 shows a comparison of the results for HuV_{NP}-IgE and MoV_{NP}-IgE in binding inhibition assays; Figure 5 shows the structure of three oligonucleotides used for site directed mutagenesis; Figure 6 shows a protocol for the construction of CDR replacements by insertion of CDR cassettes into a vector containing four framework regions fused together; Figure 7 shows the sequence of the variable domain of antibody D1.3 and the gene coding therefore; and 50 Figure 8 shows a protocol for the cloning of the D1.3 variable domain gene. This example shows the production of an altered antibody in which the variable domain of the heavy chains comprises the framework regions of a human heavy chain and the CDRs from a mouse heavy chain. The framework regions were derived from the human myeloma heavy chain NEWM, the crystallographic 55 structure of which is known (see Poljak et al., loc. cit. and Reth, M., Hammerling, G.J. and Rajewsky, K., EMBO J., 1, 629-634, 1982.) The CDRs were derived from the mouse monoclonal antibody B1-8 (see Reth et al., loc. cit.), which binds the hapten NP-cap (4-hydroxy-3-nitrophenyl acetyl-caproic acid: K_{NP-CAP}=1.2 μM). 60 A gene encoding a variable domain HuV_{NP} , comprising the B1-8 CDRs and the NEWM framework regions, was constructed by gene synthesis as follows. The amino acid sequence of the V_H domain of NEWM is shown in Figure 2, wherein it is compared to the amino acid sequence of the V_H domain of the B1-8 antibody. The sequence is divided into framework regions

and CDRs according to Kabat et al. (loc. cit.). Conserved residues are marked with a line.

The amino acid and nucleotide sequence of the HuV_{NP} gene, in which the CDRs from the B1-8 antibody

alternate with the framework regions of the NEWM antibody, is shown in Figure 3. The HuV_{NP} gene was derived by replacing sections of the MoV_{NP} gene in the vector pSV-V_{NP} (see Neuberger, M.S., Williams, G.T., Mitchell, E.B., Jouhal, S., Flanagan, J.G. and Rabbitts, T.H., Nature, 314, 268-270, 1985) by a synthetic fragment encoding the HuV_{NP} domain. Thus the 5' and 3' non-coding sequences, the leader sequence, the L-V 5 intron, five N-terminal and four C-terminal amino acids are from the MoV_{NP} gene and the rest of the coding 5 sequence is from the synthetic HuV_{NP} fragment. The oligonucleotides from which the HuV_{NP} fragment was assembled are aligned below the corresponding portion of the HuV_{NP} gene. For convenience in cloning, the ends of oligonucleotides 25 and 26b form a Hind II site followed by a Hind III site, and the sequences of the 25/26b oligonculeotides therefore differ from the 10 HuV_{NP} gene. 10 The HuV_{NPS} inthetic fragment was built as a Pstl-Hind III fragment. The nucleotide sequence was derived from the protein sequence using the computer programme ANALYSEQ (Staden, R., Nuc. Acids. Res., 12, 521-438, 1984) with optimal codon usage taken from the sequences of mouse constant domain genes. The oligonucleotides (1 to 26b, 28 in total) vary in size from 14 to 59 residues and were made on a Biosearch SAM 15 or an Applied Bio systems machine, and purified on 8M-urea polyacrylamide gels (see Sanger, F. and Coul-15 son, A., FEBS Lett., 87, 107-110, 1978). The oligonucleotides were assembled in eight single stranded blocks (A-D) containing oligonucleotides. [1,3,5,7] (Block A), [2,4,6,8] (block A'), [9,11,13a,13b] (Block B), [10a, 10b,12/14] (block B'), [15, 17] (block C), [16, 18] (block C'), [19, 21, 23, 25] (block D) and [20, 22/24, 26a, 26b) (block D'). In a typical assembly, for example of block A, 50 pmole of oligonucleotides 1,3,5 and 7 were phosphoryla-20 ted at the 5' end with T4 polynucleotide kinase and mixed together with 5 pmole of the terminal oligonucleotide [1] which had been phosphorylated with 5 μCi [γ-32p] ATP (Amersham 3000 Ci/mmole). These oligonucleotides were annealed by heating to 80°C and cooling over 30 minutes to room temperature, with unkinased oligonucleotides 2, 4 and 6 as splints, in 150 µl of 50 mM Tris.Cl, pH 7.5, 10 mM MgCl₂. For the 25 ligation, ATP (1 mM) and DTT (10mM) were added with 50 UT4 DNA ligase (Anglian Biotechnology Ltd.) and 25 incubated for 30 minutes at room temperature. EDTA was added to 10 mM, the sample was extracted with phenol, precipitated from ethanol, dissolved in 20 µl of water and boiled for 1 minute with an equal volume of formamide dyes. The sam:: lè,was loaded onto and run on a 0.3 mm 8M-urea 10% polyacrylamide gel. A band of the expected size was detacted by autoradiography and eluted by soaking. Two full length single stration were assembled from blocks A to D and A' to D' using splint oligonucleoti-30 des. Thus blocks A to D were annealed and ligated in 30 µl as set out in the previous paragraph using 100 pmole of oligonucleotides 10 t, 16 and 20 as splints. Blocks A' to D' were ligated using oligonucleotides 7, 13b and 17 as splints. After phenol/ether extraction, block A-D was annealed with block A'-D', small amounts were cloned in the 35 vector M13amp18 (Yanish-Perron, C., Vieira, J. and Messing, J., Gene, 33, 103-119, 1985) cut with Pstl and 35 Hind III, and the gene sequenced by the dideoxy technique (Sanger, F., Nicklen, S. and Coulson, A.R., PNAS USA, 74, 5463-5467, 1979). The MoV_{NP} gene was transferred as a Hind III -BamHI fragment from the vector pSV- V_{NP} (Neuberger et al., loc. cit.) to the vector M13mp8 (Messing, J. and Vieira, J., Gene, 19, 269-276, 1982). To facilitate the replace- $40\,$ ment of MoV_NP coding sequences by the synthetic HuV_NP fragment, three Hind II sites were removed from the 40 5' non-coding sequence by site directed mutagenesis, and a new Hind II site was subsequently introduced near the end of the fourth framework region (FR4 in Figure 2). By cutting the vector with Pstl and Hind II, most of the V_{NP} fragment can be inserted as a Pstl-Hind II fragment. The sequence at the Hind II site was corrected to NEWM FR4 by site directed mutagenesis. The Hind III -Bam HI fragment, now carrying the HuV_{NP} gene, was excised from M13 and cloned backinto 45 $pSV-V_{NP}$ to replace the MoV_{NP} gene and produce a vector $PSV-HuV_{NP}$. Finally, the genes for the heavy chain constant domains of human Ig E (Flanagan, J.G. and Rabbitts, T.H., EMBO J., 1, 655-660, 1982) were introduced as a Bam HI fragment to give the vector pSV-HuV_{NP}. HE. This was transfected into the myeloma line J558 L by spheroplast fusion. The sequence of the HuV_{NP} gene in pSV- HuV_{NP} . HE was checked by recloning the Hind III-Bam HI fragment 50 back into M13mp8 (Messing et al., loc. cit.). 2558L myeloma cells secrete lambda 1 light chains which have been shown to associate with heavy chains containing the MoV_{NP} variable domain to create a binding site for NP-cap or the related hapten NIP-Cap (3-iodo-4-hydroxy-5-nitrophenylacetyl-caproic acid) (Reth, M., Hammerling, G.J. and Rajewsky, K., Eur. J. Immuno!., 8, 393-400, 1978). As the plasmid pSV-HuV_{NP}.HE contains the gpt marker, stably transfected myeloma cells could be selected 55 in a medium containing mycophenolic acid. Transfectants secreted in antibody (HuV_{NP}-IgE) with heavy

chains comprising a HuV_{NP} variable domain (i.e. a "humanised" mouse variable region) and human γ con-

affinity chromatography on NIP-cap Sepharose (Sepharose is a registered trade mark). A polyacrylamide - SDS gel indicated that the protein was indistinguishable from the chimeric antibody MoV_{NP}-lgE (Neuberger

The culture supernatants of several gpt+ clones were assayed by radioimmunoassay and found to contain 60 NIP-cap binding antibody. The antibody secreted by one such clone was purified from culture supernatant by

stant domains, and lambda 1 light chains from the J558L myeloma cells.

et al., loc. cit.).

The HuV_{NP} -lgE antibody competes effectively with the MoV_{NP} -lgE for binding to both anti-human-lgE and to NIP-cap coupled to bovine serum albumin.

Various concentrations of HuV_{NP}-IgE and MoV_{NP}-IgE were used to compete the binding or radiolabelled MoV_{NP}-IgE to polyvinyl microtitre plates coated with (a) Sheep anti-human-IgE antiserum (Seward Lab-5 oratories); (b) NIP-cap-bovine serum albumin; (c) Ac38 anti-idiotypic antibody; (d) Ac 146 anti-idiotypic antibody; and (e) rabbit anti-MoV_{NP} antiserum. Binding was also carried out in the presence of MoV_{NP}-IgM antibody (Neuberger, M.S., Williams, G.T. and Fox, R.O., Nature, 312, 604-608, 1984) or of JW5/1/2 which is an lgM antibody differing from the MoV_{NP} -lgM antibody at 13 residues mainly located in the V_H CDR2 region.

The results of the binding assays are shown in Figure 4, wherein black circles represent HuV_{NP}, white 10 circles MoV_{NP}, black squares MoV_{NP}-IgM and white squares JW5/1/2. Binding is given relative to the binding in the absence of the inhibitor.

The affinities of HuV_{NP}-IgE for NP-caps and NIP-cap were then measured directly using the fluorescence quench technique and compared to those for MoVNP-IgE, using excitation at 295 nm and observing emission at 340 nm (Eisen, H.N., Methods Med. Res., 10, 115-121, 1964).

Antibody solutions were diluted at 100 nM in phosphate buffered saline filtered (0.45 μm pore cellulose acetate) and titrated with NP-cap in the range 0.2 to 20 μM. As a control, mouse DI-3 antibody (Mariuzza, R.A., Jankovic, D.L., Buot, G., Amit, A.G., Saludjian, P., Le Guern, A., Mazie, J.C. and Poljak, R.J., J. Mol. Biol., 170, 1055-1058, 1983), which does not bind hapten, was titrated in parallel.

Decrease in the ratio of the fluorescence of HuV_{NP} -lgE or HuV_{NP} -lgE to the fluorescence of the D1-3 antibody 20 was taken to be proportional to NP-cap occupancy of the antigen binding sites. The maximum quench was about 40% for both antibodies, and hapten dissociation constants were determined from least-squares fits of triplicate data sets to a hyperbola.

For NIP-cap, hapten concentration varied from 10 to 300 nM, and about 50% quenching offluorescence was observed at saturation. Since the antibody concentrations were comparable to the value of the dissocia-25 tion constants, data were fitted by least squares to an equation describing tight binding inhibition (Segal, I.H., in "Enzyme Kinetics", 73-74, Wiley, New York, 1975).

The binding constants obtained from these data for these antibodies are shown in Table 1 below.

Table 1

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	K _{NP} -cap	K _{NIP} -cap
MoV _{NP} -IgE	1.2 μM	0.02 µM
HuV _{NP} -IgE	1.9 µM	0.07 µM

These results show that the affinities of these antibodies are similar and that the change in affinity is less than would be expected for the loss of a hydrogen bond or a van der Waals contact point at the active site of an enzyme.

Thus, it has been shown that it is possible to produce an antibody specific for an artificial small hapten, comprising a variable domain having human framework regions and mouse CDRs, without any significant 40 loss of antigen binding capacity.

As shown in Figure 4(d), the HuV_{NP}-IgE antibody has lost the MoV_{NP} idiotypic determinant recognised by the antibody Ac146. Furthermore, HuV_{NP}-IgE also binds the Ac38 antibody less well (Figure 4(c)), and it is therefore not surprising the HuV_{NP}-IgE has lost many of the determinants recognised by the polyclonal rabbit anti-idiotypic antiserum (Figure 4(e)).

It can thus be seen that, although the $\text{HuV}_{\text{NP}}\text{-}\text{lgE}$ antibody has acquired substantially all the antigen binding capacity of the mouse CDRs, it has not acquired any substantial proportion of the mouse antibody's anti-

The results of Figures 4(d) and 4(e) carry a further practical implication. The mouse (or human) CDRs could be transferred from one set of human frameworks (antibody 1) to another (antibody 2). In therapy, anti-50 idiotypic antibodies generated in response to antibody 1 might well bind poorly to antibody 2. Thus, as the anti-idiotypic response starts to neutralise antibody 1 treatment could be continued with antibody 2, and the CDRs of a desired specificity used more than once.

For instance, the oligonucleotides encoding the CDRs may be used again, but with a set of oligonucleotides encoding a different set of framework regions.

The above works has shown that antigen binding characteristics can be transferred from one framework to another without loss of activity, so long as the original antibody is specific for a small hapten.

It is known that small haptens generally fit into an antigen binding cleft. However, this may not be true for natural antigens, for instance antigens comprising an epitopic site on protein or polysaccharide. For such antigens, the antibody may lack a cleft (it may only have a shallow concavity), and surface amino acid re-60 sidues may play a significant role in antigen binding. It is therefore not readily apparent that the work on artificial antigens shows conclusively that CDR replacement could be used to transfer natural antigen binding

Therefore work was carried out to see if CDR replacement could be used for this purpose. This work also involved using primer-directed, oligonucleotide site-directed mutagenesis using three synthetic oligo-65 nucleotides coding for each of the mouse CDRs and the flanking parts of framework regions to produce a

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variable domain gene similar to the HuV_{NP} gene.

Example 2

The three dimensional structure of a complex of lysozyme and the antilysozyme antibody D1.3 (Amit et al., 5 loc. cit.) was solved by X-ray crystallography. There is a large surface of interaction between the antibody and antigen. The antibody has two heavy chains of the mouse IgG1 class (H) and two Kappa light chains (K), and is denoted below as H₂K₂.

NA of

The DNA sequence of the heavy chain variable region was determined by making cDNA from the mRNA of the D1.3 hybridoma cells, and cloning into plasmid and M13 vectors. The sequence is shown in Figure 7, in which the boxed residues comprise the three CDRs and the asterisks mark residues which contact lysozyme.

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Three synthetic oligonucleotides were then designed to introduce the D1.3 V_HCDRs in place of the V_HCDRs of the HuV_{NP} gene. The Hu_{NP} gene has been cloned into M13mp8 as a BamHI-Hind III fragment, as described above. Each oligonucleotide has 12 nucleotides at the 5' end and 12 nucleotides at the 3' end which are complementary to the appropriate HuV_{NP} framework regions. The central portion of each oligonucleotide encodes either CDR1, CDR2, or CDR3 of the D1.3 antibody, as shown in Figure 5, to which reference is now made. It can be seen from this Figure that these oligonucleotides are 39, 72 and 48 nucleotides long re-

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10 pmole of D1.3 CDR1 primer was phosphorylated at the 5' end and annealed to 1µg of the M13-HuV_{NP} template and extended with the Klenow fragment of DNA polymerase in the presence of T4 DNA ligase. After 20 and oligonucleotide extension at 15°C, the sample was used to transfect E. Coli strain BHM71/18 mutLand plaques gridded and grown up as infected colonies.

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After transfer to nitrocellulose filters, the colonies were probed at room temperature with 10 pmole of D1.3 CDR1 primer labelled at the 5' end with 30 μ Ci³²-p-ATP. After a 3" wash at 60°C, autoradiography revealed about 20% of the colonies had hybridised well to the probe. All these techniques are fully described in

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25 "Oligonucleotide site-directed mutagenesis in M13" an experimental manual by P. Carter, H. Bedouelle, M.M.Y. Waye and G. Winter 1985 and published by Anglian Biotechnology Limited, Hawkins Road, Colchester, Essex CO2 8JX. Several clones were sequenced, and the replacement of HuV_{NP} CDR1 by D.13 CDR1 was confirmed. This M13 template was used in a second round of mutagenesis with D1.3 CDR2 primer; finally template with both CDRs 1&2 replaced was used in a third round of mutagenesis with D.13 CDR3 primer. In 30 this case, three rounds of mutagenesis were used.

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The variable domain containing the D1.3 CDRs was then attached to sequences encoding the heavy chain constant regions of human $\lg G2$ so as to produce a vector encoding a heavy chain Hu^* . The vector was transfected into J558L cells as above. The antibody $Hu^*_2L_2$ is secreted.

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For comparative purposes, the variable region gene for the D1.3 antibody was inserted into a suitable 35 vector and attached to a gene encoding the constant regions of mouse IgG1 to produce a gene encoding a heavy chain H* with the same sequence as H. The protocol for achieving this is shown in Figure 8.

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As shown in Figure 8, the gene encoding the D1.3 heavy chain V and $C_{\rm H}1$ domains the part of the hinge region are cloned into the M13mp9 vector.

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The vector (vector A) is then cut with Ncol, blunted with Klenow polymerase and cut with Pstl. The PStl-40 Ncol fragment is purified and cloned into Pstl-Hindll cut MV_{NP} vector to replace most of the MV_{NP} coding sequences. The MV_{NP} vector comprises the mouse variable domain gene with its promoter, 5' leader, and 5' and 3' introns cloned into M13mp9. This product is shown as vector B in Figure 8.

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Using site directed mutagenesis on the singe stranded template of vector B with two primers, the sequence encoding the N-terminal portion of the $C_H 1$ domain and the PstI site near the N-terminus of the V domain are 45 removed. Thus the V domain of D1.3 now replaces that of V_{NP} to produce vector C of Figure 8.

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Vector C is then cut with HindllI and BamHI and the fragment formed thereby is inserted into HindllI/BamHI cut M13mp9. The product is cut with Hind III and SacI and the fragment is inserted into PSV- V_{NP} cut with Hind III/SacI so as to replace the V_{NP} variable domain with the D1.3 variable domain. Mouse IgG1 constant domains are cloned into the vector as a SacI fragment to produce vector D of Figure 8.

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Vector D of Figure 8 is transfected into J558L cells and the heavy chain H* is secreted in association with the lambda light chain L as an antibody H $_2$ L $_2$.

Separated K or L light chains can be produced by treating an appropriate antibody (for instance D1.3 antibody to produce K light chains) with 2-mercaptoethanol in guanidine hydrochloride, blocking the free interchain sulphydryls with iodoacetamide and separating the dissociated heavy and light chains by HPLC in guanidine hydrochloride.

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Different heavy and light chains can be reassociated to produce functional antibodies by mixing the separated heavy and light chains, and dialysing into a non-denaturing buffer to promote re-association and refolding. Properly reassociated and folded antibody molecules can be purified on protein A-sepharose columns. Using appropriate combinations of the above procedures, the following antibodies were prepared.

7. The method of claim 6, in which the cell line which is transformed to produce the altered antibody is an

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chain respectively;

65 immortalised cell line.

c) transforming a cell line with the first or both prepared vectors; and d) culturing said transformed cell line to produce said altered antibody.

- 8. The method of claim 7, in which the immortalised cell line is a myelo na cell line or a derivative thereof.
 9. The method of any one of claims 6 to 8, in which the DNA sequence exceding the altered variable domain is prepared by oligonucleotide synthesis.
- 10. The method of any one of claims 6 to 8, in which the DNA sequence encoding the altered variable 5 domain is prepared by primer directed oligonucleotide site-directed mutagen esis using a long oligonucleotide.

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